ACYLTRANSFERASE
Inventors: David W. LEUNG et al.
Docket No.: 077319-0381



口

Phe

Leu

Leu

Leu

Leu

Phe

Leu

Leu

Leu

Leu Leu

Met

Figure 1

 Trp GGGGACAGCCCCCCCACCATTCCTACCGCTATGGGCCCAACCTCCCACTCC CACCTCCCTCCATCGGCCGGGGCTAGGACACCCCCAAATCCCGTCGCCC CCTTGGCACCGACCCCGACAGAGACAGAGACACAGCCATCCGCCACCA CCGCTGCCGCAGCCTGGCGAGGGGGCCCAGCCCCCCAGGCCCCTAC CCA GGG GCA TGG 1 GGAAGTCAGCAGGCGTTGGGGGGGGGGGGGGAATAGCGGCGGCAGC CTG CTC TTC Met Asp Leu Trp Pro Gly Ala CTG CTG TTG TGG TTCCCCTCTGAGGTGGCCAGA ATG GAT CTC CTGCTGCTG CTGATG 301 101 201 51 51 151 343 20 AAG CCC AGT GCC TGC AGC TTC CCC ACC CTG TGG 10 CIG

Lys Ala Ser Pro Ser CysPhe Trp Leu Thr Pro Leu

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Figure 1 (continued)

 \mathcal{O} 口 ATC TGG Trp G1ySGC Asn TAC AAT TyrTTCPhe 40 Ala CCC Met AAG ATG LysPhe Phe TAC 418

Val GTG Cys Ala CCC CTC GCC ATC CCT GTG TGT Pro Val Ile Val Leu Ala GTG 50 Ala TTC CTG GCT Leu Phe 455

 \mathcal{O} 口 Arg CGTLeu TTGIle ATC AAG LysATG Met Asn GAG AAC Glu GTC Val Asn AAC Arg CGC G1yGGA 9 Arg CGA 493

Arg CGA ATC G1y999 TyrTAC CTG TAC Leu TYrAAA LysIle ATC CAC His Leu CICLeu CIGMet ATG

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Figure 1 (continued)

 \mathcal{O} C Ser TCG CCC Pro Pro $T^{T}C$ Phe CAC His His GCT CAC 90 Ala G1YCGA GGG Arg Val GTG Glu GAG GTGVal 568

Leu Ser TCC AAC CAC CAG AGC TCT Ser Gln Ser Asn His GTC Val Val GTT 100 Val GTT Tyr \mathtt{TAT} Pro CCC ln 605

 \mathcal{O} H Gly Arg CGC 120 CCA GGC Pro CTGLeu GTA Val Glu GAG Met ATG GGG ATG Gly Met Leu Leu CTT110 CLGAsp GAT643

GCT GGC Trp Ala GTG CCC ATT GCC AAG CGC GAG CTA CTG TGG Leu Glu Leu Lys Arg Ala Ile ProVal γS GI 680

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Figure 1 (continued)

Д Ile ATC GGA GTC Val G1yAla CTG GCA Trp Leu 140 TGG Cys $_{
m LGC}$ CCC Gly Leu Ala CTG GGG Ala CCC Ser TCT718

Val CGG AAG CGC ACG GGG GAT GCC ATC AGT GTC Ser I1eLys Arg Thr Gly Asp Ala 150 Arg Asp GAC Ile ATC he 755

> C Gln Asp 170 GAC CAG Thr CTC ACC Gln Thr Leu Leu CIGTCT GAG GTC GCC CAG ACC Ala Glu Val Ser 160 Met ATG 793

CCT GAG GGA ACG AGA AAC CAC Glu Gly Thr Arg Asn His Pro m LLLPhe GTG Val TGG Trp GTC Val AGG 830

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180

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Figure 1 (continued)

ATG CTG CCC TTC AAA CGT GGC GCC TTC C 二 Phe Gly Ala Pro Phe Lys Arg 190 Leu Met GGC TCC Ser G1yAAT Asn 868

CCC ATA Ile Pro ATT GTC Val Ile CCC Pro AT CTT GCA GTG CAG GCC CAG GTT Gln Ala Gln Val 200 Leu Ala Val <u>1</u>8 905

G U 220 LysTGC AAG AAG LysCysTAC Tyr $T^{T}C$ Phe GAC Asp Gln CAA TAC Tyr TCC Ser Ser $_{
m LCC}$ ATG Met 210 GTCVal 943

GTG Val TCG GGA CAA TGT CAG GTG CGG Gln Val Arg CysGly Gln Ser ACC Thr TTCPhe CGC Arg CGTArg AG

230

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Figure 1 (continued)

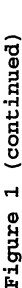
G K Asp GAT Pro ACG GAA GGG CTG ACA CCA Glu Gly Leu Thr 240 Thr CCC ProGTG Val CCA Pro CCC Pro CTGLeu 1018

GAC AGA GTC CGG CAC TCC ATG Met Ser His Arg Val Arg Asp CCA GCT CTG GCT Leu Ala 250 Ala Pro GTC Val AC 1055

U Ö TTC CGG GAA ATC TCC ACT GAT GGC CGG 270 Gly Arg Asp Thr Ser Ile Glu Phe Arg GTT Val ACT Thr 260 CTCLeu 1093

GT GGT GGT GAC TAT CTG AAG AAG CCT GGG GGC GGT GGG TGA ACCCTGGCTCTGAGCTCTCCTCCCATCTGTCCCCATCTTCCTCCC CACACCTACCCACCCAGTGGGCCCTGAAGCAGGGCCAAACCCTCTTCCTT GICTCCCCTCTCCCCACTTATTCTCCTCTTTGGAATCTTCAACTTCTGAA ly Gly Gly Asp Tyr Leu Lys Lys Pro Gly Gly Gly 280 1130 1266 1168

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ACTCTTGCCTCGGTGCAGTTTCCACTCTTGACCCCCCACCTCCTACTGTCT TGTCTGTGGGACAGTTGCCTCCCCCTCATCTCCAGTGACTCAGCCTACAC CTCTACCCCTCTACCCCCACATTGGCCCAGTGGACTCATCCATTCTTTGGA AAGGGAGGGAACATTCCATCCCCAGTGGAGTCTCTTCCTATGTGGTCTT ACAAATCCCCCCCCCTCCAAAGTCCATGGATTCAATGGACTCATTT TGTGAGGAGGACTTCTCGCCCTCTGGCTGGAAGCTGATACCTGAAGCACT TGTAGCCTCCTGTCAGTGGGGGCTGGACCCTTCTAATTCAGAGGTCTCAT GCCTGCCCTTGCCCAGATGCCCAGGGTCGTGCACTCTCTGGGATACCAGT TCAGTCTCCACATTTCTGGTTTTCTGTCCCCATAGTACAGTTCTTCAGTG GACATGACCCCACCCAGCCCCTGCAGCCCTGCTGACCATCTCACCAGAC ACAAGGGGAAGAAGCAGACATCAGGTGCTGCACTCACTTCTGCCCCCTGG GGAGTTGGGGAAAGGAACGAACCCTGGCTGGAGGGGATAGGAGGGCTTTT 1316 1466 995. 1366 1416 1516 616 999. 1766 1816 1716 1966 1866 1916

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Figure 1 (continued)

CCCACACAGGGGGGGGGCTGGGAGCAAAAGGAGGGGTGGGACCCAG 2016 AATTTATTTCTTTTCTGTTGAGGCTTCCCCCTCTCTGAGCCAGTTTTCA TTTCTTCCTGGTGGCATTAGCCACTCCCTGCTCTCACTCCAGACCTGTT TTTTGCGTGGTTGTTTTTTTTATTATTATCTGGATAACAGCAAAAAACTG 2066 2116 2166

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2216 AAAATAAAGAGAGAGAGAAAAAAAA

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50 <u>F</u> <u>F</u> KMA G <u>F</u> VTIRP <u>F</u> SKS	100 -YGIRVEVRG MLGLDVK -FGLKVECRK WAGVKVQLHA	150 -VPI-AKRELVTAKKSL TVTV-GKKSL STLAVMKKSS	LTQDVRVWV- KKNKRALWV- KKRRISIWM- KDFPRPFWLA	250 LAVQAQVPIV LAQQGKIPIV AAIAAGVPII SAVSIMRDFY
40 <u>LWFCSPSAKY</u> C ILVCVFGSIY <u>LIV</u> NAIQAVL	90 RLMLLHIKYL ARCFYHVMKL GHMFGRLAPL WLQLVWVVDW	140 VLPGRC IFPPGCT IVQ-PP ILAQRSGCLG	SVMSEVAQTL SVMSEVAQTL DTLNKGLENV GTIAEVVNHF KTLKWGLQRL	240 MLPFKRGAFH MLPFKKGAFH -LPFKTGAFH VLIPRTKGFV
30 LL-LLFLLPT AL-AG RL-IITVIYS PLGLLFLLSG	80 GRNVENMKIL GKQHLAQWIT PRNPKHVATF	130 SLDLLGMME STLDIFMLGR NNYDMVTASN SDIDWL-IGW	180 IDRKRTGDAI LDRSKROEAI IDRNNRTKAH LERS-WAKDE	230 T T ASQGLPAPRN
20 <u>LLL</u> LLF YLRSV <u>L</u> - <u>VVL</u> <u>LYIF</u> <u>PL</u> V <u>LVV</u> L	70 VLAIPVCAVR VIASILCTLI CLFS EL	120 -PYVVSNHQ KPYIMIANHQ -NAIYIANHQ EHALIISNHR	170 WLAGVIE WEMALSGTYE WLTGNLL WFAEYLE	220 GS SEL RGL
10 <u>M</u> DLWPGAWM- <u>M</u> SV-IGRFLY <u>M</u>	60 FYNGWILFLA FYGG FYRINRFLA	110 AHHF-PPSQ- VVGE-ENLAK PTDA-ESYG- DEETYRSMGK	160 <u>LW</u> AGSAGLAC <u>KYVPFLG</u> <u>LWIPFFGQLY</u> <u>KFLPVIGWSM</u>	210 -FPEGTRNHN -FPEGTRSYT -FPEGTRSRG LFVEGTRFTP
ਜਿਜਜ	51 51 51 51	101 101 101 101	151 151 151 151	201 201 201 201
Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT



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Inventors: David W. LEUNG et
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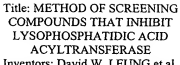
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290 300 VPTEGLTPDD VPALADR ISTENLTKDK IGEFAEK IDVSQYGKDQ VRELAAH VIHVRMKRHA MSEMPKSDED	340 GDYLKKPGGG G* DTTLPPQ	390 400	430 440 450
280 GQCQVRVLPP GCMIVRILKP GLVIVEMLPP MLRILKGQSS	330 REISTDGRGG KEIGYSPAIN AELDKEVAE- ATGTFDEEIR	380 HDKKVNKKIK FTAAGMALVT	430
270 YCKKERRFTS VSPKYGVFNR INLNRLHN IVPKDSPQPT	320 VRHSMLTV-F VRDQMVDT-L CRSIMEQK-I AKDALLDKHL	370 A <u>L</u> Q Q <u>L</u> LSTWRGVA	420 KKMH*
260 PIVMSSYQDF PVVVSNTSTL PVCVSTTSNK PALYDTTV	310 VSKWCKDIFV	360 <u>AIE</u> YA FG <u>AIE</u> FFKWT	410 <u>s</u> <u>v</u>
251 251 251 251	301 301 301 301	351 351 351 351	401 401 401 401
Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT	Human LPAAT Yeast LPAAT E.coli LPAAT Maize LPAAT

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60	120	180	240	300	360	420	480	540
GCCGTGTCTG	CGAGTTCTAC	CTCGCTCGTC	CTGGTTCGTG	CAGGCTGCAĠ	GATGGGCCTC	CTTCCTGGGG	GCGCTCTAGC	CCTCAAAGTG
50	110	170	230	290	330 340 350 360 cercreaac caccagacca recressions	400 410	470	530
TGGAGCTGTG	GCCGCGCGC	CCGCCGTGGC	GCATCATCGG	GGGACCCGCG		ATCGCCAAGC GGGAGCTGCT	TCAACCGGCA	TCAGGGAGAA
40	100	160	220	280	340	400	450 · 460	520
GGCCGGGCCA	GTGCAGCTGA	TTCACGGTGT	GAGAACATGA	TTCGAGGTGC	CACCAGAGCA	ATCGCCAAGC	CCTCGGGGG GTCTTCTTCA	GAGCGCATGG
20 30	90	150	210	270	330	390	450	510
GGCGCGCCG TCGGGCGCCG	GCTGCTGCTG	CGCGCTGTGC	CCGGACGGTG	CGGGCTCCGC	CGTCTCCAAC	CTGCGTGCAG	CCTCGGGGGC	CGACCTGGGC
	80	130	200	260	310	380	440	500
	TGCTGTTGCT	GCCAAGGTCG CCCTGTACTG	GCCACGGCGG	AGTACTTTTA	GAGGCCCGTC CCTGTGTCAT	TTCCGGAGCG	TCATCATGTA	CAGTGATGGC
10 GGAGCGAGCT	0 <i>L</i>	130 GCCAAGGTCG	190 TGCCTGCTGT	250 CGAAGCTTCA	310 GAGGCCCGTC	370 ATGGAGGTCC	430 CCCGTGGGCC	490 530 540 540 ACTGCCATGA CAGTGATGG CGACCTGGGC GAGCGCATGG TCAGGGAGAA CCTCAAAGTG



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Figure 3 (continued)

600	660	720	780	840	900	960	1020	1080
Taagaagggg	CTCTTCCTTC	AGTGCAGGTG	CGTGGACACC	GGAGAACGGG	GGGCATGACC	CTCCCGGCTT	AGGAAGCCCC	CAGCTGGACC
590	650	710	770	820 830	890	950	1010	1070
TGCTGCCTTT	CCGTGGTGTA	GAACAGTCAC	TCCCTGCGCT	CACATCTCCA AGACCCCCA	GACCACGGCA	GAGGGGACTC	CCCGGGAAGC	GCAGGGGGCT
580	640	680 690 700 710 720 ACAACACCAA GAAGAAGTTC TTCACTTCAG GAACAGTCAC AGTGCAGGTG	760	820	880	940	1000	1060
AATGGGGACC	CCCATCGTCC		GCGGCGGACG	CACATCTCCA	CAGTAGCCCA	AGGATGGGCA	TCTCACTCAG	GGTGTCCCCT
570	630	690	750	810	870	930	990	1050
TCGCAACGAC	GGCACAGGTG	GAAGAAGTTC	CGGCCTCACT	CACCTTCCTC	GCAGCCGGCC	CGATGGCTGG	CTCCCCAGC	ACAGGCCCCT
560	620		740	800	860	910 920 930 940	980	1040
CCGAGGGTAC	TGGCAGTCCA		TCCCCACCAG	CCATGAGGAC	GGTCTGGCGT	TGGGGAGGGC AGGTGGAAGC CGATGGCTGG AGGATGGGCA	CTCTGTCCGG	GGTCTCAGAC
550	610	670	730	790	850	910	970	1030
TGGATCTATC	GCCTTCTACC	TCCTCCTTCT	CTGGAAGCCA	TGCCACCGGG	GCCACTGCGG	TGGGGAGGGC	CCAAATACCA	TTCTGTCACT

Figure 3 (continued)

04	04	O (1)	O 75	O 4	_
1140	1200	1260	1320	1380	1440
GGGNTGATAA	CGATGGCCCA	GGGCCACAGG	GTCAGCACTG	AAAAAAAAA	
1130	1190	1250	1310	1370	1430
TCTGGGNGCT	ACAÄGGCCCC	CACGCACCCT	AATCTGTGGG	ААААААААА	
1190 1100 1110 1120 1130 1140	1150 1160 1170 1180 1190 1200	1210 1220 1230 1240 1250 1260	1270 1280 1290 1300 1310 1320	1330 1340 1350 1360 1370 1380	1420
CTCCCGGGC TCGAGGGCAG GGACTCGCGC CCACGGCACC TCTGGGNGCT GGGNTGATAA	AGATGAGGCT TGCGGCTGTG GGGCTGAGCC ACAÄGGCCCC CGATGGCCCA	GGAGCAGATG GGAGGACCCCAGGA GTCCCAGACT CACGCACCT GGGCCACAGG	GAGCCGGGAA TCGGGGCCTG CTGCTCCTGC TGGCCTGAAG AATCTGTGGG GTCAGCACTG	TACTCCGTTG CTGTTTTTT ATAAACACAC TCTTGGAAAA AAAAAAAAAA	
1110	1170	1230	. 1290	1350	1410
GGACTCGCGC	GCCCGCTGGT	GAGGCCAGGA	CTGCTCCTGC	ATAAACACAC	
1100	1160	1220	1280	1340	1400
TCGAGGGCAG	TGCGGCTGTG	GGAGGACCCC	TCGGGGCCTG	CTGTTTTTT	
1090	1150	1210	1270	1330	1390
CTCCCGGGC	AGATGAGGCT	GGAGCAGATG	GAGCCGGGAA	TACTCCGTTG	AAA

Figure 4

CCG 50 TGG Trp Leu CTGGAG Glu Met GGAGCGAGCTGGCGCCGTCGGGCGCCCGGGCCCGGGCC ATG 30 20

CTG Leu 20 CAG Gln GTG Val 90 CTG Leu Leu CTGCIGLeu CTGLen Leu TTGLeu CTG Ala Leu GCG CTG Ala gcg CCC CTGLen TGT Cys

GCG CysTAC TyrLeu CTG GTC GCC Val Ala 120 TAC GCC AAG (Lys Ala TYrPhe GAG TTC Glu 225 525 Ala Ala Ser AGC 100

Leu 50 CTG CTGCys Leu $_{
m TGC}$ 180 GTC Val Leu CIC Ser TCG GTG GCC Ala 170 Val gcc Ala Ser GTG TCC 160 Val TTC Phe 150 TGC CTG Leu

TGG Trp ATC GGC Ile AAC ATG AGC ATC Ser Asn Met 60 GAG Glu 210 GTG ACG CGG Arg GGC Gly Gly 200 ggc TGC Cys 190

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		GTG	Val 80		GTC	Val		TC	Val 110		TTC	Phe		5 S	Phe	140		GAC	Asp
																7			
		GAG	Glu	320	ATC	Ile		GAG	Glu	410	CTG CTC	Leu		TTC	Phe		200	ATG GCC	Ala
		TTC	Phe		GTC	Val		ATG	Met	•	CTG	Leu		GTC	Val		υ,	ATG	Met Ala
	270	CGC	Arg		TGT	Cys	360	CTC	Leu		GAG	Glu	450	ggc	G1y			GTG	Val
		CTC	Leu	310	CGT CCC	Pro		GGC		0	CGG	Arg		GGG	G1y		0	ACA	Thr
nuec		GGG	G1y	31	CGT	Arg 90		ATG	Met	400	AAG CGG	Lys Arg 120		CTC	Leu		490	ATG ACA	Met 150
(continued	260	TAC	Phe Tyr (Ala	350	GAC ATG	Asp Met		CCC	Ala	440	TAC	Tyr			gcc	Ala
		$ ext{T} ext{L}$	Phe		GAG	Glu	(-)				ATC	Ile	7	ATG TAC	Met				\mathtt{Thr}
ce 4		TAC	$\Gamma Y \mathcal{X}$	300	CAG	Gln			Leu	390	CAG	Gln		ATC	Ile		480	AGC	Ser
Figure 4	250	TTC AAG	Lys		CTG	Leu	340	AGC ATC	Ile		GTG	Val	430	CTC	Leu			TCT	Ser
124	2				AGG	Arg	ñ				TGC	Cys	43	9	Gly Leu	130		CGC	Arg
		AGC	Ser	290	ධවට වටට	Arg		CAG		380	GAG CGC	Arg		GTG	Val		470	CGG CAG	Gln
			Arg			Pro		CAC	His	(-)	GAG	Glu		သည	Pro		7	CGG	Arg
	240	GTG	Val		GAC	Asp	330	AAC	Asn		SSS	Pro	420	GGG	G1y			AAC	Asn
		TTC	Phe	280	990	Arg		TCC	Ser	370	CT T	Leu		CTG	Leu		460	ATC	Ile

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Figure 4 (continued)

•	$\begin{array}{c} \mathtt{TAT} \\ \mathtt{TY} \\ \mathtt{170} \end{array}$	AAG Lys	GTC Val 200	AAG Lys	GCC Ala 230
	ATC Ile	590 TTT Phe	ATC Ile	680 AAG Lys	GAA
	TGG	CCT (CCC	680 ACC AAG Thr Lys	CTG
	540 GTG Val	CTG	630 GTG Val	AAC Asn	720 GTG Val
	aaa Lys	CTG CTG Leu	CAG Gln		CAG Gln
	CTC	580 GAC CTG ASP Leu 180	GCA Ala	670 TTC TAC Phe Tyr 210	GTG Val
	530 GAG AAC Glu Asn	GGG Gly	620 GTC CAG Val Gln	TCC Ser	710 GTC ACA (Val Thr
	GAG Glu	AAT Asn	6 GTC Val	TCC	7 GTC Val
	agg Arg	570 GAC ASP	GCA Ala	660 TTC Phe	CA
	520 ATG GTC Met Val 160	AAC Asn	610 TAC CTG TYr Leu 190	TCT TCC Ser Ser	0 GGA 31Y
	52 ATG Met 160	CGC Arg	610 TAC CT TYr Le 190	TCT Ser	700 TCA G(Ser G] 220
	CGC Arg	560 ACT Thr	TTC Phe	650 GTG TAC Val Tyr	ACT Thr
	GAG Glu	GGT A	GCC Ala	GTG Val	TTC Phe
	510 GGC Gly	GAG Glu	600 GGC Gly	GTG Val	690 TTC Phe
	CTG	550 CCC Pro	AAG Lys	640 CCC Pro	AAG Lys



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Figure 4	

	GTG		
770	CTC	Leu	
	gce	Ala	
	CCT GCG CTC	Pro	
0.0	GTC	Asp Val B	
760	GAC	Asp	240
	GCG	Ala	
	SCG	Ala	
750	ACT	Thr	
	CTC	Leu	
•	ggc	Gly	
740	AGC	Ser	
1-	ACC	Thr	
	222	Pro	
730	ATC	Ile	

	TCC	Ser	260
	ATC		
	CAC	His	
810	CTC	Leu	
	TTC	Phe	
	ACC	Thr	
800	ACC	Thr	
ω	AGG	Arg	
	ATG	Met	
790	ပ္ပင္ပ		
7.5	CGG	Arg	250
	CAC	His	
	TGC	Cys	
780	ACC	Thr	
	GAC	Asp	

	CAG	Gln		
860	GTG	Val		
ω	399	G1y		
	TCT	Ser		
850	9999	G1y		
8	gcg	Ala	270	
	ACT	Thr		
	gcc	Ala		
840	GGG	Gly		
	AAC	Asn		
	GAG	Glu		
830	CAG	Gln		
ω	CC	Pro	*	
	ACC	Thr	•	
820	AAG	Lys		
Φ				

	870			880	068	006	910	920
CCG	CCC	CAG	TAG	CCCAGACCA	CGGCAGGGCA	NTGACCTGGG	CCG GCC CAG TAG CCCAGACCACGGCAGGGCATGACCTGGGGAGGGCAGGTGGAAGC	GAAGC
Pro	Pro Ala Gln ***	Gln	* * *					

	CTCAGAC	1100	AGGGCAG
0	CTCCCCCAGCTCTCACTCAGCCCGGGAAGCAGGAAGCCCCCTTCTGTCACTGGTCTCAGAC	1090	ACAGGCCCCTGGTGTCCCCTGCAGGGGGCTCAGCTGGACCCTCCCCGGGCTCGAGGGCAG
1040	GCCCCTTC	1080	GGACCCTC
1030	AGCAGGAA	1070	GCTCAGCT
1020	AGCCCGGG2	1060	CTGCAGGGG
1010	TCTCACTC		GGTGTCCC
1000	CTCCCCCAGO	1050	ACAGGCCCCT

Title: METHOD OF SCREENING COMPOUNDS THAT INHIBIT LYSOPHOSPHATIDIC ACID

ACYLTRANSFERASE Inventors: David W. LEUNG et al. Docket No.: 077319-0381

Figure 4 (continued)

1110	1120	1130	1140	1150	1120 1130 1140 1150 1160
	GGACTCGC	GCCCACGGCA	CCTCTGGGNG	CTGGGNTGATA	GGACTCGCCCCACGGCACCTCTGGGNGCTGGGNTGATAAAGATGAGGCTTGCGGCTGTG
1170	1180	1190	1200	1210	1180 1190 1200 1210 1220
	GCCCGCTG	GTGGCTGAG	SCACAAGGCCC	ccgarggccc	GCCCGCTGGTGGCCTGAGGCCCCCCGATGGCCCAGGAGCAGGAGGACCCC
1230	1240	1250	1260	1270	1240 1250 1260 1270 1280
	GAGGCCAG	GAGTCCCAGA(STCACGCACC	TGGGCCACAG	GAGGCCAGGAGTCCCAGACTCACGCACCTGGGCCACAGGGAGGCCGGGAATCGGGGCCTG
1290	1300	1310	1320	1330	1300 1310 1320 1330 1340
	CTGCTCCT	GCTGGCCTGAA	AGAATCTGTGG	GGTCAGCACT	CTGCTCCTGCTGACAAAATCTGTGGGGTCAGCACTGTACTCCGTTGCTGTTTTTT
1350	1360 ATAAACAC	1360 1370 1380 ATAAACACTCTTGGAAAAAAAAAAAAAAAAAAA	1380 \AAAAAAAAA	AAAAAAAA	aaaa





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Figure 5

Alignment of LPAAT Sequences.

10 20 40 50 1	1 MAKTRUSS-L RNRRQLKP AVAATADD DKDGVFMV 1 MDASCASSFL RGRCLESCFK ASFGMSQPKD AAGQPSRRPA DADDFFTVDD	51 QLSRAE FYAKVAL YC ALG TVSAVA SLVCLLGGG RIVENM-SII 51 QLSRAE FYAKVAL YC ALG TVSAVA SLVCLLGGG RIVENM-SII 51 TIWFCS BAK YFFKWAF-YN GWILFIAVIA IPVCAV-RG RNVENM-KIL	51	51 <u>MLYI FRLIV'IVIX'S ILVCVFG SLICLIDEM FANIN</u> 51LLSC FKIFVCFAFT VVLITAVAWG LIMVILLIDEWP YMRIRLGNLY	51 DRWITVILSV VRIPACEL- SMMTIIWN MIMILLINEWE YAKIEVEN
Human LPAAT-β Human LPAAT-α Yeast LPAAT E.coli LPAAT H.influenzae	S.typnimuriu L.douglassi C. nucifera	Human LPAAT-β Human LPAAT-α	.Yeast LPAAl' E.coli LPAAT H.influenzae	S.typhimuriu L.douglassi	C. nucifera

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Figure 5 (continued)

140 150	GWEVRSFKYFYGLRFEV RDPRRLQEAR PCVIVSNHQS ILDMGIMEV RLMITHIKYLYGIRVEV RGAHFFPSQ PYVVSNHQS SLDILGMEV CFY-HVMKLMLGLDVKY VGEENLAK-K PYIMIANHQS TLDIEMGRI GHMFGRL APLFGLKVEC RKPTDAESYG NAIYIANHQN NYDMVTASNI ARWFGRL -FT YPLFGLKVEH RIPQDQKQIS RAIYIGNHQN NYDMVTASNI GHMFGRL -FT APLFGLKVEC RKPADAENYG NAIYIANHQN NYDMVTAANI GHIGGLV - IMIYGIPIKI QGSEHIKKRA IFTYISNHAS PIDAFFVMU GHUGRLY IMIYGIPIKI QGSEHIKKRA IFTYISNHAS PIDAFFVMU GHUGRLY MILGNPITI EGSEFSNIRA I YICNHAS IVDIFIIMM	160 170 180 190 200 LPERCYQLAK RELLFIGEV GLIMYLGGV FETINECRSST AMTVMAIL LPGRCV PIAK RELLMAGSA GLACWIAGV IFIDRIKRIGD ALS - VMSEV VQPPTVTVGK KSLIMIPFF GQLYWLIGN ILIDRININRIK AHGTIAEV VQPPTVTVGK KSLIMIPFF TGLIYWVIGN ILIDRININRIK AHGTIAEV VQPPTVTVGK KSLIMIPFF TGLIYWVIGN ILIDRININRIK AHGTIAAV VQPPTVTVGK KSLIMIPFF TGQLYWLIGN ILIDRININRIK AHGTIAAV 1 PKGTVT IAK KEVI WYPILG QLYTLAH IRIDRSN PAA ALGSFTMKEA
130	RDPRRLQEAR E RGAHHFPPSQ I VGEENLAK-K E VGEENLAK-K E RYPTDAESYG RYPTDAESYG RYPTDAESYG RYPTDAESYG RYPADAENYG GSEHIKKRA I EGSEFSNITRA	180 -GLIMYLGGV -GLYCWLAGV -GWFWALSGT -GOLYWLTGN TGILYWVIGN TGILYWVIGN TGILYWVIGN QPTLYVLAHH QPTLYVLAHH
120	FYGLRFEVLYGIRVEVMLGLDVKV APLFGLKVE YPLFGLKVEH APLFGLKVEH IMIYGIPIKI MMILGNPITI	
110	101 GWEVRSF KY – 101 RLMITHI KY – 101 CFY – HVM KL – 101 CHMFGRL – — 101 CHMFGRL – FT	151 <u>IP ERCY QIAK</u> 151 <u>IP ERCY PIAK</u> 151 <u>IP GRCY PIAK</u> 151 <u>VQP PIVITY GK</u> 151 <u>VQP PIVITY GK</u> 151 <u>VQP PIVITY GK</u> 151 <u>VQP PIVITY GK</u> 151 <u>IP IG TY GVAK</u> 151 1 <u>P IG TY GVAK</u> 151 1 <u>P IG TY GVAK</u>
	Human LPAAT-β Human LPAAT-α Yeast LPAAT E.coli LPAAT H.influenzae S.typhimuriu L.douglassi C. nucifera	Human LPAAT-β Human LPAAT-α Yeast LPAAT E.coli LPAAT H.influenzae S.typhimuriu L.douglassi C. nucifera

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continued)
Figure 5 (0

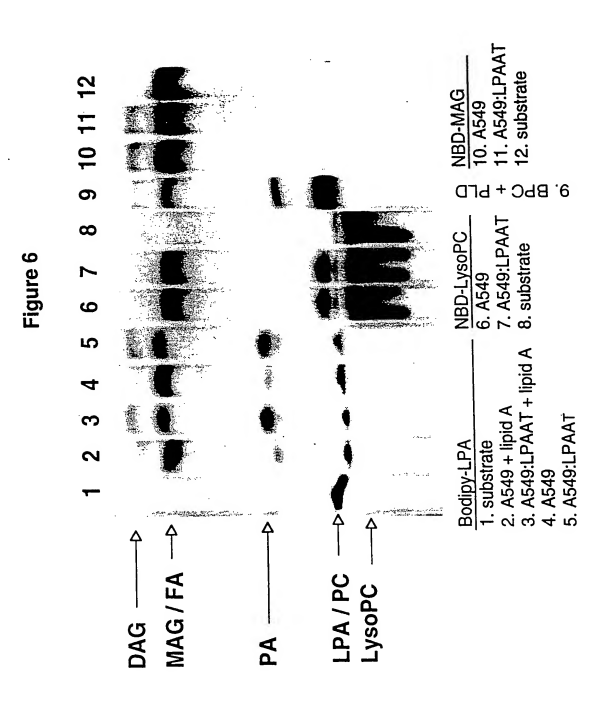
210 GERMARENLK VMI YPEGTRN DNGDL—LPF KKGAFTL—A VQAQVPIVPV 201 AQIILIQDAR VWVFPEGTRN HNGSM—LPF KRGAFTL—A VQAQVPIVP I 201 LENVKKNRRA LWVFPEGTRS YISELIMLPF KKGAFTL—A QQGALPIVPV 201 VNHFKKRRS S IWMFPEGTRS RGRGL—LPF KTGAF—HAA LAGVPLIPV 201 VNHFKKRRS S IWMFPEGTRS RGRGL—LPF KTGAF—HAA LAGVPLIPV 201 VNHFKKRRS S IWMFPEGTRS RGRGL—LPF KTGAF—HAA LAGVPLIPV 201 VNHFKKRRS S IWMFPEGTRS GJGRL—LPF KTGAF—HAA LAGVPLIPV 201 VRVITHKNIS LIMFPEGTRS GJGRL—LPF KTGFHHAA LAGSH_PIVPM 201 VRVITHKNIS LIMFPEGTRS GJGRL—LPF KTGFHHFITA LQIRL_PIVPM 201 VRVITHKNIS LIMFPEGTRS KTGRL—LPF KTGFHHFITA LQIRL_PIVPM	251 VYSSFSS—FYNTKKFFTS GTVTVQVLEA IPTSGLTAD VPALVDTCHR 251 VMSSYQD—F YCKK RRFTS GQQQYRVLPP VPT GGLTPDD VPALADRVRH 251 VWSNTST—L VSPKYGVFNR GMIVRILKP ISTENLITKDK ICEFFAEKVRD 251 VVSNTST—— NK INLARIHN GLVTVEMLPP IDVSGYGKDQ VRELAAHCR— 251 VCSSTH—— NK INLARINN GLVTVEMLPP IDVSGYTKDN VRDLAA YCHF 251 VCSSTH—— NK INLARINN GLVTVEMLPP VDVSGYGKDQ VRELAAHCRE 251 VCSSTH—— NK INLARINN GLVTVEMLPP VDVSGYGKDQ VRELAAHCRE 251 VCSSTH—— NK INLARINN GLVTVEMLPP INTDDMTVDK IDDYV KMIHD 251 LITGITHLAWF TRKGIFRVRP VPITVKYLPP INTDDMEEK INTYVEMIHE
Human LPAAT-α Yeast LPAAT E.coli LPAAT H.influenzae S.typhimuriu L.douglassi C. nucifera	Human LPAAT-β Human LPAAT-α Yeast LPAAT E.coli LPAAT H.influenzae S.typhimuriu L.douglassi

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Figure 5 (continued)

		310	320	330	340	350
Human LPAAT- β	301	AMRTIFIHIS		GSGVQPAQ*-		
Human LPAAT-α	301	SMLTVFREIS	LDGROGGDVL KKPGOGG*-	KKPGGGG*		
Yeast LPAAT	301	OMVDIL KEIG	OMVDILKEIG YSPAINDTTE PROAIEYAAL OHDKKVNKKI KNEPVPSVSI	PPQAIEYAAL	QHDKKWNKKI	KNEPVPSVSI
E.coli L.PAAT	301	-SIMEOKIAE	LDKEVA — ER	EAAGKV*		
H.influenzae	301	TILMEKRIAE	LD EETA	KGN *	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
S. typhimuriu	301	TALMEOKIAE	INIMEOKIAE LDKEVA — ER EATCKV*	EATCKV*	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1
L.douglassi	301	IYVRNLPASQ	IYVRNILPASQ KPLGSTINRS-K*-	-S-K*		
C. nucifera	301	TALYVOHLPE	$\overline{\mathbf{I}}$ ALYVDHLPE SQ $\overline{\mathbf{K}}$ PLVSKG $\overline{\mathbf{R}}$ D $\overline{\mathbf{ASG}}$ RS $\overline{\mathbf{N}}$ S*	ASGESNS*-		
		360	370	380	390	
Human LPAAT-β	351		•	•	•	•
Human L.PAAT- α	351			•		•
Yeast LPAAT	351	SINDWITHNEG	SSVKKMH*	•	•	•
E.coli L.PAAT	351			•	•	•
H.influenzae	351			•	•	•
S. typhimuriu	351	 - - - -		•	•	•
L.douglassi	351		•		•	•
C. nucifera	351		•	•	•	•

Docket No.: 077319-0381



TLC Analysis of Acyltransferase Acitvity

Docket No.: 077319-0381

Induction of TNF in A549 LPAAT or A549 cells stimulated with mTNF and IL-1

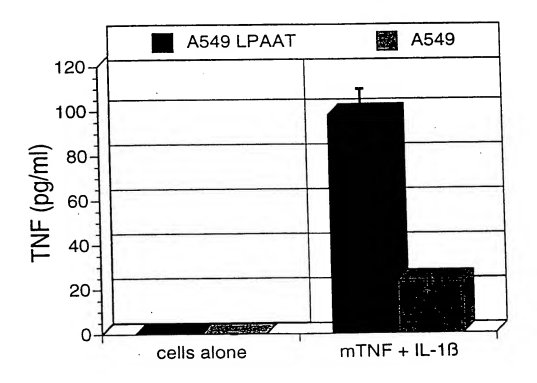


Figure 7

Docket No.: 077319-0381

Induction of IL-6 in A549 LPAAT or A549 cells stimulated with mTNF and IL-1

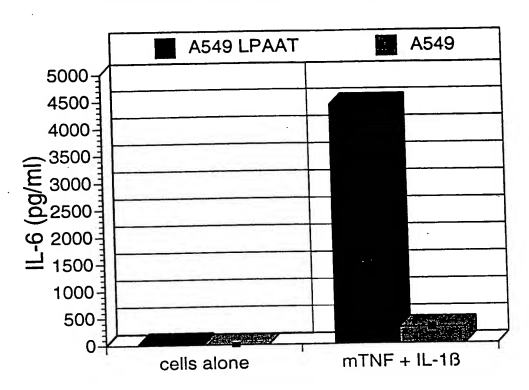


Figure 8

Inventors: David W. LEUNG et al. Docket No.: 077319-0381

Figure 9 Translated sequence of human LPAAT-γl

GGCĀ	GGTC	TCTG	GCTT	GTCC	ACCC(GGAA(CATCA GCCCT AGCGA	GAG GGGG	GCA(GCCG	TGCA	CCCG	CTCC	TGAG	CAGC	60 120 180
TGAC	CTTC	1666	CMC	ACGG.		בכוכו	TG A	AG	ACC	CAG	TTC (GTG	CTG	CAC	225
GCC	<u>ATG</u>	GGC	CIG	CIG	31-	Dho I	Leu I	ve '	Thr (Gln	Phe '	Va l	Leu	His	
	Met	GIY	Leu	Leu .		Pne .	Jeu i	3 y 3		10					
					5 ama 1	mm <i>c</i> (-mc (TC	ACT .		CTG .	GTC	Δ'nC	AAC	270
CTG	CTG	GTC	GGC	TTT	GTC .	TTC	GTG (71G .	Co.	C112	LOU	Val	Tla	Acn	
Leu	Leu	Val	Gly	Phe	Val	Pne	Val V	/aı	ser	25	neu	Val	116	A311	
15					20				mcc.		CITIC	NCC	770	CAC	315
TTC	GTC	CAG	CTG	TGC	ACG	CTG	GCG (TIC	TGG	CCG	G1C	AGC	AAG	CAG	213
Phe	Val	Gln	Leu	Cys	Thr	Leu .	Ala	_eu	rrp	Pro	vaı	ser	Lys	GIII	
2.0					35					4.0					360
CTC	TAC	CGC	CGC	CTC	AAC	TGC	CGC (CTC	GCA	TAC	TCA	CTC	166	AGC	360
Leu	Tvr	Arg	Arg	Leu	Asn	Cys	Arg	Leu	Ala	Tyr	Ser	Leu	Trp	Ser	
4 =					50					22					405
	CTG	GTC	ATG	CTG	CTG	GAG	TGG '	rgg	TCC	TGC	ACG	GAG	TGT	ACA	405
Gln	T.eu	Val	Met	Leu	Leu	Glu	Trp	Trp	Ser	Cys	Thr	Glu	Cys	Thr	
					65					70					
	ጥጥር	ACG	GAC	CAG	GCC	ACG	GTA	GAG	CGC	TTT	GGG	AAG	GAG	CAC	450
Lou	Dhe	Thr	Asp	Gln	Ala	Thr	Val	Glu	Arg	Phe	Gly	Lys	Glu	His	
					80					82					
75	CEC	አ ጥር	ልጥር	СТС	AAC	CAC	AAC	TTC	GAG	ATC	GAC	TTC	CTC	TGT	495
GCA	GTC	TIO	TIO	LAU	Asn	His	Asn	Phe	Glu	Ile	Asp	Phe	Leu	Cys	
	Val	TTG	TIE	Пеп	95				3	.00	_			_	
90			3 mc	mcm.	GAG	CGC	TTC	GGA	GTG	CTG	GGG	AGC	TCC	AAG	540
GGG	TGG	ACC	AIG	161	Clu	724	Phe	Glv	Va1	Leu	Glv	Ser	Ser	Lvs	
		Thr	Mec	Cys	110	ALG	£ 11C	011	•	115	1				
105					110	CTC	CTC	ጥልሮ	GTG		CTC	ATC	GGC	TGG	585
GTC	CTC	GCT	AAG	AAG	GAG	CIG	CIC	TAC	Val	Pro	Leu	Tle	Glv	Tro	
Val	Leu	Ala	Lys	Lys	GIU	Leu	Leu	TAT	vai	130	Deu		013		
120					125	3 mm	CEC.	mmc	ሞርር		CGG	AAC	TCC	GAG	630
ACG	TGG	TAC	TTT	CTG	GAG	ATT	GTG	Dho	Cre	Luc	7~~	Tare	Trn	Glu	
Thr	Trp	Tyr	Phe	Leu	Glu	TIE	Val	Pne	Cys	145	Arg	пуз	110	014	
135	i				140			000	cmc		ccc	CTC	TCG	GAC	675
GAG	GAC	: CGG	' GAC	ACC	GTG	GTC	GAA	666	CIG	AGG	3~~	tou	502	Acn	3,3
Glu	Asp	Arg	Asp	Thr	Val	Val	Glu	GIY	rea	160	Arg	Leu	261	νaδ	
					755					TOO					720
TAC	: cc	GAG	TAC	ATG	TGG	TTT	CTC	CTG	TAC	TGC	GAG	000	ACG	CGC	720
Tyr	Pro	Glu	Tyr	Met	Trp	Phe	Leu	Leu	TYT	Cys	GIU	GIY	THE	Arg	
	-				170					T/2					765
		GAG	ACC	AAG	CAC	CGC	GTT	AGC	ATG	GAG	GIG	GCG	GCL	GCT	765
Phe	Th	Glu	Thr	Lys	His	Arg	Val	Ser	Met	GIU	vai	Ala	. Ala	Ala	
40/	•				185					720	,				010
		G CTI	CCI	GTC	CTC	AAG	TAC	CAC	CTG	CTG	CCG	CGG	ACC	AAG	810
Live	Gly	r Leu	. Pro	Val	Leu	Lys	Tyr	His	Leu	. ьеч	Pro	Arg	Thr	Lys	
	-				200					200)				
		~ ACC	ACC	GCA	GTC	AAG	TGC	CTC	CGG	GGG	ACA	GTC	: GCA	GCT	855
Cl	r Dh	e Thi	Thr	Ala	Val	Lys	Cys	Leu	Arg	, Gly	Thr	: Val	. Ala	Ala	
	_				215					22	,				
21		m C 7 1	ኮ ርጥ፤	ACC	י כידכי	AAC	TTC	AGA	GGA	AAC	AAC	AAC	ccc	TCC	900
GT	TA	I GA.	. 17-1	The	. T.air	Asn	Phe	Arc	Gly	Ası	ı Lys	s Asr	n Pro	Ser	
	_				つてい	1				43.	,				
22	5 		~ am/	- CTC	ጉ ጥልር		AAG	AAC	TAC	GAC	G GCC	GAG	TATO	TGC	945
CT	GCT	اقاق اقا مام	J ATC		, m	. GGG	Tave	Live	יע"ו	Gli	ı Ala	a Ast	o Me	Cys	
		u GI	A TIE	e ner	TINE	. <u>G</u> ly	د بر	~J ~	3 -	250)			•	
24	0				245	,	GNO	<u>አ</u> ጥር	י ככל			י קא	A A A	G GAA	990
GT	G AG	G AG	A TT	r cci	r CTC	GAA	A OAC	יות.	. Dec	n Ter	1 Acr	יום מ	1 LV	G GAA	
		g Ar	g Phe	e Pro	o re	ı Gil	LASP	116		26				s Glu	
25	5				260) 		m x /	י כאי	20. 20.	- 774	G G M	- 60	3 CTC	1035
GC	A GC	T CA	G TG	G CT	r CA	r AAA	r C.1.0	TA	_ CA	י הי	~ AA	- GA		G CTC	

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Figure 9 (continued)

Ala	Ala	Gln	Trp	Leu	His	Lys	Leu	Tyr	Gln	Glu 280	Lys	qzA	Ala	Leu	
270 CAG Gln	GAG Glu	ATA Ile	TAT Tyr	AAT Asn	CAG Gln	AAG Lys	GGC Gly	ATG Met	TTT Phe	CCA Pro 295	GGG Gly	GAG Glu	CAG Gln	TTT Phe	1080
285 AAG Lys					CCG Pro	maa	3.00	CTC	CTG	AAC	TTC	CTG	TCC Ser	TGG	1125
300 GCC Ala					TCT Ser	666	CTC.	ጥጥር	AGT	TTT	GTC	TTG	GGC Gly	GTC	1170
315 TTT Phe					CCT Pro		CITIC	አጥር	CTG	ACT	TTC	TTG	GGG	TTT Phe	1215
330 GTG Val	GGA Gly	GCA Ala	GCT Ala	TCC Ser	TTT Phe	GGA Gly	GTT Val	CGC Arg	AGA Arg	CTG Leu	ATA Ile	GGA Gly	GTA Val	ACT Thr	1260
345 GAG Glu	ATA	GAA Glu	AAA Lys	GGC Gly	, Ser	AGC Ser	TAC	GGA Gly	AAC Asn	CAA	GAG	TTT Phe	Lys	AAA Lys	1305
360	GAA	. TAA	TTF	ATG	365 CTGT	GACT	GAAC	ACAC	:GCGG	CCCT	rGAC	GTGC	TATO	CAGTT	1362
Lys AAC CTC TG!	Glu TCA GTG AAGT(AACC ACTA! CTTC!	CAAC! ATAT! AGCC!	ACACA LOAA1 LOCCA	AGAGT AAAAC ACAGC	rgcac cttg <i>i</i>	GAAA AGCCA GGT(AAGAC AAGAC CCCAC	CAATT TAAT CATC	TAGAA AGAA CTCCA	AACTI TTCA(ACGC(GAAC)	ATTTT SAAGO SCGCO TCCGO	TTCTT GCCT(CCGT(CATTAA GTCAGG GGAGG CAAGAG	1422 1482 1542 1602 1660

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Figure 10 Translated sequence of LPAAT-72 cDNA

CACGCTGGCGCTCTGGCCGGTCAGCAAGCAGCTCTACCGCCGCCTCAACTGCCGCCTCGCC TACTCACTCTGGAGCCTAGCACAAAACTAGAAGCAACCCAAGCACCTGTCACTGGAGACT 121 AATTATGCGGCACCCATACAGGGACCCTCTGCGGCCATCATGGAGAGCCTTCATCTTGCC 181 CGTACAGTTTTAAGCGAAAAAAGGAAGTATACAACAAAGTCCATAACTGGTC ATG CTG 238															
CGT	ACAG'	rtt <u>t</u>	AAGC	GAAA	AAGG2	AAGT	ATAC	AACA	AAGT	CCA <u>T</u>	AACT	GGTC		CTG Leu	238
				TCC											283
		5		Ser			10					15	-		
GCC	ACG	GTA	GAG	CGC	TTT	GGG	AAG	GAG	CAC	GCA	GTC	ATC	ATC	CTC	328
		20		Arg			25					30			
AAC	CAC	AAC	TTC	GAG	ATC	GAC	TTC	CTC	TGT	GGG	TGG	ACC	ATG	TGT	373
		35		Glu			40					45		_	
				GTG											418
		50		Val			55					60	-	-	
				GTG											463
		65		Val			70				-	75			
				TGC											508
		80		Cys			85				-	90	-		
GTG	GTC	GAA	GGG	CTG Leu	AGG	CGC	CTG	TCG	GAC	TAC	CCC	GAG	TAC	ATG	553
		95					100					105	_		
				TAC Tyr											598
-		110					115					120		_	
				ATG Met											643
		125					130			_	_	135	•		400
				CTG Leu											688
	_	140		CGG			145			-		150			722
				Arg											733
	_	155		GGA			160				_	165			778
				Gly											//0
		170	_	TAC		_	175					180			823
				Tyr											023
-1-	1	185	-1-	-1 -			190		-1-		3	195		113	
				CCG											868
•		200		Pro			205					210	_		
				CAG											913
	Ī	215	_	Gln			220					225	_		
				TTT											958
	_	230		Phe			235					240	_		
				CTG											1003
	-	245		Leu			250		_			255			
				AGT											1048
Ser	Pro	Leu	Phe	Ser	Phe	Val	Leu	Gly	Val	Phe	Ala	Ser	Gly	Ser	

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Figure 10 (continued)

		260					265					270			
ССТ	CTC	CTG	ATC	CTG	ACT	TTC	TTG	GGG	TTT	GTG	GGA	GCA	GCT	TCC	1093
Pro	Leu	Leu	Ile	Leu	Thr	Phe	Leu	Gly	Phe	Val	Gly	Ala	Ala	Ser	
		275					280					285			
ተመተ	GGA	GTT	CGC	AGA	CTG	ATA	GGA	GTA	ACT	GAG	ATA	GAA	AAA	GGC	1138
Phe	Glv	Val	Arg	Arq	Leu	Ile	Gly	Val	Thr	Glu	Ile	Glu	Lys	Gly	
	_	290					295					300			
TCC	AGC	TAC	GGA	AAC	CAA	GAG	TTT	AAG	AAA	AAG	GAA	TAA	TTA	ATGGC	1185
Ser	Ser	Tvr	Gly	Asn	Gln	Glu	Phe	Lys	Lys	Lys	Glu	***			
		305					310								
тсто	TACT(SAAC	ACAC	GCGG	CCT	SACG	GTGG'	TATC	CAGT	raac'	rcaa.	AACC	AACA	CACAG	1245
A CTO	CACC	מממב	CAC	ል ልጥጥ	GAA	ACTA	$\mathbf{T}\mathbf{T}\mathbf{T}\mathbf{T}$	TCTT?	ATTA	ACTG	GTGA(CTAA'	ratt.	AACAA	1305
770	ኮጥር እ	CCA	AGAG'	TAAA	TAAT	rcag.	AAGG	CCTGʻ	rcag	GTGA	AGTC'	TTCA(GCCT	CCCAC	1365
N.C.C	~~ > ~ (COTO	CAG	ר אתרי	TCCAC	CGCG	CGCC	CGTG	ggag(GTGG	GTCC	GGCC(GGAG.	AGGCC	1425
TCC	CGCG	GACG	CCGT	CTCT	CCAG	AACT	CCGC'	TTCC	AAGA	GGGA	CCTT'	TGGC'	TGCT'	TTCTC	1485
TCC'	TTAA	ACTT.	AGAT	CAAA	TTTT	AAAA	AAAA	AAAA	AAA						1523

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Figure 11 Translated sequence of human LPAAT- δ

GAAGGAGACGCCTTCCTGAGTCCTGGATCTTTCTTCCTTC													61 121 175		
									Met	Asp	Leu	Ala	G1y 5	Leu	1.5
CTG	AAG	TCT	CAG '	TTC (CTG	TGC	CAC (CTG	GTC	TTC	TGC	TAC	GTC	TTT	220
Leu	Lys	Ser	Gln :	Phe	Leu	Cys	His !	Leu 15	Val	Phe	Cys	Tyr	Val 20	Phe	
ATT	GCC	TCA	GGG (CTA .	ATC	ATC	AAC .	ACC	ATT	CAG	CTC	TTC	ACT	CTC	265
Ile	Ala	Ser	Gly :	Leu	Ile	Ile	Asn '	Thr 30	Ile	Gln	Leu	Phe	Thr 35	Leu	244
CTC	CTC	TGG	CCC .	TTA	AAC	AAG	CAG	CTC	TTC	CGG	AAG	ATC	AAC	TGC	310
		Trp	40					45					50		255
AGA	CTG	TCC	TAT	TGC	ATC	TCA	AGC	CAG	CTG	GTG	ATG	CTG	CTG	GAG	355
		Ser	55					60					65		400
TGG	TGG	TCG	GGC	ACG	GAA	TGC	ACC	ATC	TTC	ACG	GAC	CCG	200	Ala	400
_		Ser	70					75					80		445
TAC	CTC	AAG Lys	TAT	GGG	AAG	GAA	AAT	81a	TIO	Val	Ual	Lau	AAC	Hie	447
_		Lys GAA	25					90					95		490
AAG	TTT	GAA Glu	ATT	GAC	L.L.I.	LOU	CAS	Glv	Trn	Ser	Leu	Ser	Glu	Ara	450
			100					105					TTO		
ጥጥጥ	GGG	CTG	ጥጥ እ	GGG	GGC	TCC	AAG	GTC	CTG	GCC	AAG	AAA	GAG	CTG	535
Phe	Gly	Leu	Leu	Gly	Gly	Ser	Lys	Val 120	Leu	Ala	Lys	Lys	125	Leu	
GCC	ጥልጥ	GTC	CCA	ATT	ATC	GGC	TGG	ATG	TGG	TAC	TTC	ACC	GAG	ATG	580
Ala	Tyr	Val	Pro	Ile	Ile	Gly	Trp	Met 135	Trp	TYT	Pne	Thr	140	Met	
GTC	TTC	TGT	TCG	CGC	AAG	TGG	GAG	CAG	GAT	CGC	AAG	ACG	GTT	GCC	625
Val	Phe	Cys	Ser	Arg	Lys	Trp	Glu	GIn 150	Asp	Arg	Lys	Thr	Val 155	Ala	670
ACC	AGT	TTG	CAG	CAC	CTC	CGG	GAC	TAC	CCC	GAG	AAG	TAT	T.L.I.	Dho	670
			160					165					1/0		715
CTC	TTA	CAC	TGT	GAG	GGC	ACA	724	Dha	Thr	GAG	TAG	Live	Hic	GAG Glu	113
			175					TRO					TAS)	
ልጥር	AGC	ATG	CAG	GTG	GCC	CGG	GCC	AAG	GGG	CTG	CCT	CGC	CTC	AAG	760
Ile	. Ser	Met	Gln	Val	Ala	Arg	Ala	Lys	GIY	' Leu	Pro	Arg	Let	ı Lys	
			100					195					200	}	805
CAT	CAC	: CTG	TTG	CCA	CGA	ACC Thr	LVS	Glv	. IIC	Ala	Tle	Thr	. Ual	AGG L Arg	003
			205					210)				21	•	850
AGO	C TTC	3 AGA	AAT	G'I'A	GIT	COX	. GCI	Val	יניתו לעד	Asr	CV	Thi	Lei	AAT Asn	030
			220					225)				230	J	895
TT	C AGA	TAA A	' AAT	GAA	, AA'I	CCA	ACA	to	, [0.	, Gl	r Wal	LIZ	, Ac.	GGA	0,5
			235					240	J				24:		
λλι	2 22	מיד ב	י כאיד	CCA	GAT	TTC	TAT	GTI	r AGO	G AGO	TA E	CCZ	A CT	G GAA	940
Ly	s Ly	s Tyr	His	Ala	Asp	Leu	ı Tyr	259	L Arq	g Aro	116	e Pro	о Lei	u Giu	
GA	C ATO	c cci	מ מס י	GAC	GAT	GAC	GAG	TG	C TC	G GC	TG	G CT	G CA	C AAG	985
As	p Il	e Pro	Glu 265	. Asp	Asp	Asp	Glu	270	s Se	r Ala	a Trp	, Le	u Hi 27	ѕ шуѕ	



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Figure 11 (continued)

Leu Tyr Gln Glu Lys Asp Ala Phe Gln Glu Glu Tyr Tyr Arg Thr									030	CAC	CAC	ጥልር	ጥልር	ACC	ACG	1030
GGC ACC TTC CCA GAG ACG CCC ATG GTG CCC CCG CGG CGC TGG 1075 Gly Thr Phe Pro Glu Thr Pro Met Val Pro Pro Arg Arg Pro Trp 295 ACC CTC GTG AAC TGG CTG TTT TGG GCC TCG CTG GTG CTC TAC CCT 1120 Thr Leu Val Asn Trp Leu Phe Trp Ala Ser Leu Val Leu Tyr Pro 310 TTC TTC CAG TTC CTG GTC AGC ATG ATG ATC AGG AGC GGG TCT TCC CTG 1165 Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Ser Leu 325 ACG CTG GCC AGC TTC ATC CTC GTC TTC TTT GTG GCC TCG GTG GGA 1210 Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 345 GTT CGA TGG ATG ATT GGT GTG AGC GAA ATT GAC AAG GGC TCT GCC 1255 Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 366 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG 1301 Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGAACCCTTGGGGAACTGGTGGGCACGGCGGAAGTCACGACCTCT 1421 CCAGCCAAGGGAGTCTGGTCTCAAGGCCCGGATGGGGAAGATGTTTTGTAATCTTTTTT TCCCCATGTGCTTTTAGTGGGGCTTTTGGTTTTCTTTTTTTT	CTC	TAC	CAG	GAG	AAG	GAT	GCC	1-1-1	CAG	GAG	GAG	TAC	Tree	7.2	Thr	1030
GGC ACC TTC CCA GAG ACG CCC ATG GTG CCC CCC CGG CGG CCC TGG Gly Thr Phe Pro Glu Thr Pro Met Val Pro Pro Arg Arg Pro Trp 295 ACC CTC GTG AAC TGG CTG TTT TGG GCC TCG CTG GTG CTC TAC CCT Thr Leu Val Asn Trp Leu Phe Trp Ala Ser Leu Val Leu Tyr Pro 310 TTC TTC CAG TTC CTG GTC AGC ATG ATG ATG AGG AGG GGG TCT TCC CTG Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Ser Leu 325 ACG CTG GCC AGC TTC ATC CTC GTC TTC TTT GTG GCC TCG GTG GGA Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 356 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG TYC Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGCACCTTGGGGAACTGGGTAGGCACGGCGGAAGTCACGACCTCT 1481 1541	Leu	Tyr	Gln		Lys	Asp	Ala	Phe	GIN	GIU	GIU	TAT	IYL	200	1111	
Carrell				280							~~~		ccc		TCC	1075
ACC CTC GTG AAC TGG CTG TTT TGG GCC TCG CTG GTG CTC TAC CCT Thr Leu Val Asn Trp Leu Phe Trp Ala Ser Leu Val Leu Tyr Pro	GGC	ACC	TTC	CCA	GAG	ACG	CCC	ATG	GTG	CCC	CCC	200	200	200	166	1012
ACC CTC GTG AAC TGG CTG TTT TGG GCC TCG CTG GTG CTC TAC CCT Thr Leu Val Asn Trp Leu Phe Trp Ala Ser Leu Val Leu Tyr Pro	Gly	Thr	Phe		Glu	Thr	Pro	Met	Val	Pro	Pro	arg	Arg	PIO	Trp	
Thr Leu Val Asn Trp Leu Phe Trp Ala Ser Leu Val Leu Tyr Pro 310 TTC TTC CAG TTC CTG GTC AGC ATG ATC AGG AGC GGG TCT TCC CTG Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Ser Leu 325 ACG CTG GCC AGC TTC ATC CTC GTC TTC TTT GTG GCC TCC GTG GGA Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG TYr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT GCCCATGTGCCTTTAGTGGGCTTTTGTTTTTTTTTT				295												1100
TTC TTC CAG TTC CTG GTC AGC ATG ATC AGG AGC GGG TCT TCC CTG Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Ser Leu 325 ACG CTG GCC AGC TTC ATC CTC GTC TTC TTT GTG GCC TCC GTG GGA Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT GGGCACACGGGAGTCTGGTCTCAAGGCCGGATGGGGAAGAAGATGTTTTTTTT	ACC	CTC	GTG	AAC	TGG	CTG	TTT	TGG	GCC	TCG	CTG	GTG	CTC	TAC		1120
TTC CAG TTC CTG GTC AGC ATC AGG AGC GGG TCT TCC CTG 1165 Phe Phe Leu Val Ser Met Ile Arg Ser Gly Ser Leu 335 ACG CTG GCC AGC TTC ATC CTC GTC TTT TTT GTG GCC TCC GGA 1210 Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val GIy Ser Val ATT GAC AAC GAC AAC ATT GAC AAC ATT GAC AAT GAC AAT GAC AAG AAT GAC AAC AAT GAC AAC AAC AAA CTG AAA CTG AAA CTG AAA CTG AAA AAA CTG AAA AAA	Thr	Leu	Val	Asn	Trp	Leu	Phe	Trp	Ala	Ser	Leu	Val	Leu	Tyr	Pro	
Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Leu 325 330 335 335 335 325 325 1210 ACG CTG GCC ATC CTC GTC TTT GTT GTG GGC TCC GTG GGA Phe Phe Val Ala Ser Val Gly 350 350 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1256 1256 1256				310												
Phe Phe Gln Phe Leu Val Ser Met Ile Arg Ser Gly Ser Leu 325 330 335 335 335 325 325 1210 ACG CTG GCC ATC CTC GTC TTT GTT GTG GGC TCC GTG GGA Phe Phe Val Ala Ser Val Gly 350 350 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1255 1256 1256 1256	TTC	TTC	CAG	TTC	CTG	GTC	AGC	ATG	ATC	AGG	AGC	GGG	TCT	TCC	CTG	1165
325 ACG CTG GCC AGC TTC ATC CTC GTC TTC TTT GTG GCC TCC GTG GGA Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT GGGACACGGTGACAAAGGCTGGTGAGCCCCTTGCTGGGCACGGCGGAAGTCACGACCTCT CCCCCATGTGCCTTTAGTGGGCTTTTCTTTT	Phe	Phe	Gln	Phe	Leu	Val	Ser	Met	Ile	Arg	Ser	Gly	Ser	Ser	Leu	
Thr Leu Ala Ser Phe Ile Leu Val Phe Phe Val Ala Ser Val Gly 340 345 350 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 360 365 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG 1301 Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT 1361 GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT 1421 CCCCCATGTGCTTTAGTGGGCTTTTGTTTTTTTTTTTT				325					330					335		
340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT GGGACACGGTGACAAAGGCTGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCCACCCATGTGCTTTAGTGGGCTTTTCTTTT	ACG	CTG	GCC	AGC	TTC	ATC	CTC	GTC	TTC	TTT	GTG	GCC	TCC	GTG	GGA	1210
340 GTT CGA TGG ATG ATT GGT GTG ACG GAA ATT GAC AAG GGC TCT GCC Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT GGGACACGGTGACAAAGGCTGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCCACCCATGTGCTTTAGTGGGCTTTTCTTTT	Thr	Leu	Ala	Ser	Phe	Ile	Leu	Val	Phe	Phe	Val	Ala	Ser	Val	Gly	
Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 360 365 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG 1301 Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCCACCCATGTGCTTTAGTGGGCTTTCTTTTTTTTTT				340					345					350		
Val Arg Trp Met Ile Gly Val Thr Glu Ile Asp Lys Gly Ser Ala 355 360 365 TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG 1301 Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCCACCCATGTGCTTTAGTGGGCTTTCTTTTTTTTTT	GTT	CGA	TGG	ATG	ATT	GGT	GTG	ACG	GAA	ATT	GAC	AAG	GGC	TCT	GCC	
TAC GGC AAC TCT GAC AGC AAG CAG AAA CTG AAT GAC TGA CTCAGGG Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAAGATGTTTTTTTT	Val	Arg	Trp	Met	Ile	Gly	Val	Thr	Glu	Ile	Asp	Lys	Gly	Ser	Ala	366
Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCAAGTCACGACCTCT CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAGAAGATGTTTTTTTT				355					360					365		
Tyr Gly Asn Ser Asp Ser Lys Gln Lys Leu Asn Asp *** 370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT GGGACACGGTGACAAAGGCTGGGTGACCCCTGCTGGGCACGGCAAGTCACGACCTCT CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAAGATGTTTTTTTT	TAC	GGC	AAC	TCT	GAC	AGC	AAG	CAG	AAA	CTG	AAT	GAC	TGA	CTC	AGGG	1301
370 375 AGGTGTCACCATCCGAAGGGAACCTTGGGGAACTGGTGGCCTCTGCATATCCTCCTTAGT 1361 GGGACACGGTGACAAAGGCTGGGTGACCCCTGCTGGGCACGGCAAGTCACGACCTCT 1421 CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAGAAGATGTTTTGTAATCTTTTTT 1481 TCCCCATGTGCTTTAGTGGGCTTTGGTTTTCTTTTTTTTGTGCGAGTGTGTGAGAATGGCT 1541	Tvr	Glv	Asn	Ser	Asp	Ser	Lys	Gln	Lys	Leu	Asn	Asp	***			
GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAGAAGATGTTTTGTAATCTTTTTT TCCCCATGTGCTTTAGTGGGCTTTGGTTTTCTTTTTTTTT	_	_		370					375							
GGGACACGGTGACAAAGGCTGGGTGAGCCCCTGCTGGGCACGGCGGAAGTCACGACCTCT CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAGAAGATGTTTTGTAATCTTTTTT TCCCCATGTGCTTTAGTGGGCTTTGGTTTTCTTTTTTTTT	AGGT	GTCA	CCAT	CCGA	AGGG	AACC	TTGG	GGAA	CTGG	TGGC	CTCT	GCAT	'ATCC	TCCT	TAGT	
CCAGCCAGGGAGTCTGGTCTCAAGGCCGGATGGGGAGGAAGATGTTTTTTTT	CCCA	CACG	CTCA	CAAA	GGCT	GGGT	GAGC	CCCT	GCTG	GGCA	.CGGC	GGAA	GTCA	CGAC	CTCT	
TCCCC ATCTCCTTTAGTGGGCTTTGGTTTTCTTTTTGTGCGAGTGTGTGAGAATGGCT 1541	CCAC	CCAG	CCAC	ጥርጥር	CTCT	CAAG	GCCG	GATG	GGGA	.GGAA	GATG	TTTT	GTAA	TCTT	"I"I"I"I	
	TOCO	ር አጥር	ጥርርጥ	ጥጥልር	TGGG	CTTT	GGTT	TTCT	$\mathbf{T}\mathbf{T}\mathbf{T}\mathbf{T}$	GTGC	GAGT	'GTGI	GTGA	GAAI	GGCT	
GTGTGGTGAGTGTGAACTTTGTTGTGTGATCATAGAAAGGGTATTTTAGGCTGCAGGGGA 1601	CTCT	COTO	АСТС	TGAA	CTTT	GTTC	TGTG	ATCA	TAGA	$\mathbf{A}\mathbf{A}\mathbf{G}\mathbf{G}$	GTAT	TTTP	GGCI	GCAG	GGGA	
GCCCAGGCCTCGGGACCGAAGGGGACAAGTTCCCCTTTCATCCTTTGGTGCTGAGTTTTC 1661	cccc	ACCC	CTCC	GGAC	CGAA	GGGG	ACAA	GTTC	CCCI	'TTÇA	$^{'}$ TCCI	TTGG	TGCI	GAGI	TTTC	
TITA ACCOUTTGCTTGCCAGAGATAAAGTGAAAAAGTGCTTTAGGTGAGATGACTAAATTAT 1/21	ጥርጥል	ACCC	ጥጥርር	TTGC	CAGA	GATA	AAGT	GAAA	AGTG	CTTI	AGGI	GAGA	TGAC	TAAA	TATT	
GCCTCCAAGAAAAAAATTAAAGTGCTTTTCTGGGTCAAAAAAAA	GCCT	CCAA	GAAA	AAAA	AATI	'AAAC	TGCT	TTTC	TGGG	TCAA		AAA	LAAA	\A		1774

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Figure 12

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	10	20	30	40	50
	MGLLAFLKTQ	FVT.HLLVGFV	FVVSGLVINF	VQ-LCTLALW	PVSKQLYRRL
LPAAT-γ1					
LPAAT-Y2	MDLAGLLKSQ	THE CULTUE CVAL	FTASGLITNT	IO-LFTLLLW	PINKQLFRKI
LPAAT-δ	MDLAGLLKSQ	FUCHTARCIA	1 11.002		
	60	70	80	90	100
	NODE AVELUE	QLVMLLEWWS	CTECTLFTDQ	ATVERFGKEH	AVIILNHNFE
LPAAT-Y1		MAT I ETABLIS	CTECTLETDO	ATVERFGREH	VALITINGME
LPAAT-Y2		QLVMLLEWWS	CTECTTETOP	RANTKYGKEN	AIVVLNHKFE
LPAAT-δ	NCRLSYCISS	OFAWT FEMMS	GILCILI		
	110	120	130	140	150
		ERFGVLGSSK	VLAKKELLYV	PLIGWTWYFL	EIVFCKRKWE
LPAAT-γ1		PRESENT CESY	MAKKELLYV	PLIGWTWYFL	EIALCKKKME
LPAAT-γ2	IDFLCGWIMC	ERFGVLGSSK	ALVKKELVAN	PITGWMWYFT	EMVFCSRKWE
LPAAT- δ	IDFLCGWSES	ERFGLLGGSK	VERKKEDATV	2110///	
	160	170	180	190	200
		nnt CDVPEVM	WFLLYCEGTR	FTETKHRVSM	EVAAAKGLPV
LPAAT-γ1		CDVDEVM	WELLYCEGTR	FTFTKHRVSM	EVAAAKGLPV
LPAAT-Y2	EDRDTVVEGL	RRLSDIPEIM	TEL THOEGRE	FTEKKHEISM	QVARAKGLPR
LPAAT- δ	QDRKTVATSL	GHTKDAREK :	FFEINCEGIN	I I I I I I I I I I I I I I I I I I I	
	210	220	230		
		CEMMATAKCI'S	GTVAAVYDVI	LNF-RGNKNP	SLLGILYGKK
LPAAT-Y1		CEMMANNCT.B	CTVA AVYDVI	'LNF-RGNKNE	SPEGIFICAN
LPAAT-γ2	PKAHPPbk.rk	GFTTAVRCHR	MARSAUVDCT	I.NF-RNNENE	T LLGVLNGKK
LPAAT-δ	LKHHLLPRTK	(GFALTVRSLE	NVVSAVIBEI		₹
	260	270	280		
_		e prentet.DEA	C EAAOWLHKL	QEKDALQEIY	NOKEMERCEO
LPAAT-Y1		- STEDIDIDES	z FAAOWIHKU	Y OEKDALOEL:	NOKOME PGEQ
LPAAT-Y2	YEADMCVRR	- Predicado	ECSAWI.HKI.	Y OEKDAFOEE	YRTGTFPETP
LPAAT- δ	YHADLYVRR.	I ALEDIAEDDI) LCGL		
	310	n 329	33	0 34	
_		T NET CMATT	L LSPLFSFVL	G VFASGSPLL	ILTFLGFV
LPAAT-Yl		TOWNS CLINDS	r rodiesevi.	G VEASGSPLL	T P.L.E. P.G.E.A.
LPAAT-Y2	FKPARRPWT	P PULTOWELL	U LVPEFOFLV	S MTRSGSSLT	LASFILVF
LPAAT- δ	MVPPRRPWT	r AMMPEMWani	V DILLEGEDV	0	_
	36	ი 37	0 38	0	
	0.2	L IGVTEIEKG		K KKE*	
LPAAT-Y1	GAASFGVRR	L IGVTEIEKG	S SYGNOEF	K KKE*	
LPAAT-γ2	GAASFGVRR	T IGALETEKO	S PAGNEDERU	K LND*	
LPAAT- δ	FVASVGVRW	M IGVTEIDKG	'S MIGHODOM		



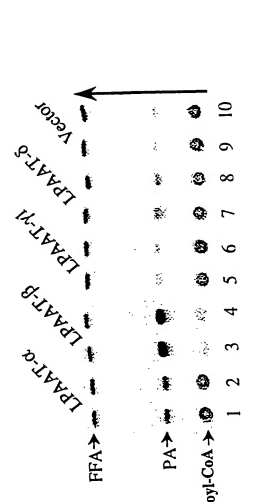


Figure 13